

CH2 CH2 CH2 CH2 CH2 \rightarrow CH2 @14 15 16 @17 @23 @24

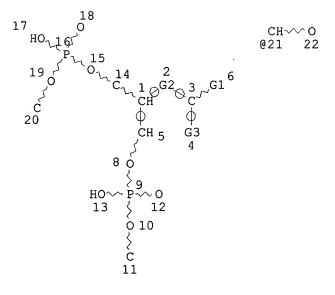
Str. Claims 5\$10\$

CH2~CH2~O~CH2~CH2 @18 19 20 21 @22

VAR G1=14-5 17-7/23-5 24-7/18-5 22-7 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 24

STEREO ATTRIBUTES: NONE L7 STR



(A= C) and 21-

VAR G1=H/HY VAR G2=N/O/P/S VAR G3=CH2/21 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

**BEST AVAILABLE COPY** 

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 21

Searcher :

Shears

308-4994

STEREO ATTRIBUTES: NONE

13:43 SEA FILE=REGISTRY SSS FUL L2 OR L7

100.0% PROCESSED 42747 ITERATIONS

SEARCH TIME: 00.00.23

1143 ANSWERS

(FILE 'HCAPLUS' ENTERED AT 15:30:13 ON 09 OCT 2002)

L10 383 S L9

L11 20 S L10 AND (?MICROB? OR ?BACTER?)

E1 THROUGH E52 ASSIGNED

L11 ANSWER 1 OF 20 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2002:696696 HCAPLUS

TITLE: Charge tags and the separation of nucleic acid

molecules

INVENTOR(S): Lyamichev, Victor; Skrzpczynski, Zbigniew;

Allawi, Hatim T.; Wayland, Sarah R.; Takova,

Tsetska; Neri, Bruce P.

PATENT ASSIGNEE(S): Third Wave Technologies, Inc., USA

Patent

SOURCE: U.S. Pat. Appl. Publ., 120 pp., Cont.-in-part of

U. S. Ser. No. 333,145.

CODEN: USXXCO

DOCUMENT TYPE:

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 11

PATENT INFORMATION:

PATENT	NO.	KIND	DATE		APPL	DATE							
US 6001	 128465 567 063030	A 19991214			US 20 US 19 WO 20								
₩0 2002 ₩:	AE, AG, CN, CO,	AL, AM CR, CU	, AT, AU, , CZ, DE,	AZ, DK,	BA, BB, DM, DZ,	BG, BF	, BY	, BZ, , FI,	CA, GB,	GD,			
	LC, LK,	LR, LS	, HU, ID, , LT, LU, , PL, PT,	LV,	MA, MD,	MG, ME	, MN	, MW,	MX,	MZ,			
	TM, TN,	TR, TT	, TZ, UA, , MD, RU,	ŪĠ,	US, UZ,		•		•	•			
RW:	CH, CY,	DE, DK	, MW, MZ, , ES, FI, , CF, CG,	FR,	GB, GR,	IE, IT	, LU	, MC,	NL,	PT,			
PRIORITY APP	SN, TD,	TG	,,,	US 1996-682853					A2 19960712				
US 1999-333145 A2 19990614 US 1996-599491 A2 19960124 US 2001-777430 A 20010206													

AB The present invention provides charge tags for attachment to materials including solid supports and nucleic acids, wherein the charge tags increase or decrease the net charge of the material. Thus, when an oligonucleotide modified with a charge tag is reduced in size (cleaved) or increased in size (elongated), the resulting product bears a net charge or a charge to mass ratio different from the original oligonucleotide thereby permitting sepn. of the original and product oligonucleotides on the basis of charge. The present invention therefore further provides methods for sepg. and

characterizing mols. based on the charge differentials between modified and unmodified materials, e.g., by capillary electrophoresis. Thus, MCP1 and ubiquitin transcripts were simultaneously detected in an in vitro assay using the Invader technol. and probes charge tagged with one of two phosphoramidates, i.e., dG-P-Cy3 or dC-P-Cy3 in which P = -O-P(:O) (NHCH2CH2NMe2)O-. ΙT 446017-73-8D, oligonucleotide conjugates 446017-74-9D, oligonucleotide conjugates RL: ARU (Analytical role, unclassified); PRP (Properties); ANST (Analytical study) (charge tags and sepn. of nucleic acid mols.) 446017-73-8 HCAPLUS RN 3'-Uridylic acid, 5-[3-[(2-aminoethyl)amino]-3-oxo-1-propenyl]-2'-deoxy-5'-O-[[3-[2,3-dihydro-2-[3-[1-(3-hydroxypropyl)-3,3-dimethyl-CN 3H-indolium-2-yl]-2-propenylidene]-3,3-dimethyl-1H-indol-1yl]propoxy]hydroxyphosphinyl]uridylyl-(3'.fwdarw.5')-5-[3-[(2aminoethyl)amino]-3-oxo-1-propenyl]-2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

H2N Me Me Me Me Me OH OH OH OH OH SENERGE 1-A

PAGE 1-B

PAGE 2-A H203P0

RN

446017-74-9 HCAPLUS
3'-Uridylic acid, 5-[3-[(6-aminohexyl)amino]-3-oxo-1-propenyl]-2'-deoxy-5'-O-[[3-[2,3-dihydro-2-[3-[1-(3-hydroxypropyl)-3,3-dimethyl-3H-indolium-2-yl]-2-propenylidene]-3,3-dimethyl-1H-indol-1-CN yl]propoxy]hydroxyphosphinyl]uridylyl-(3'.fwdarw.5')-5-[3-[(6-aminohexyl)amino]-3-oxo-1-propenyl]-2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

> Shears 308-4994 Searcher :

PAGE 1-A

PAGE 1-B

PAGE 2-B

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(CH<sub>2</sub>) 6
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L11 ANSWER 2 OF 20 HCAPLUS COPYRIGHT 2002 ACS 2002:615883 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 137:164653

Charge tags and separation of nucleic acid TITLE:

molecules

INVENTOR(S): Lyamichev, Victor; Skrzpczynski, Zbigniew;

Allawi, Hatim T.; Wayland, Sarah R.; Takova,

Tsetska; Neri, Bruce P.

Third Wave Technologies, Inc., USA PATENT ASSIGNEE(S):

PCT Int. Appl., 197 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

```
APPLICATION NO.
           PATENT NO.
                                                KIND DATE
                                                                                           WO 2002-US3423
           WO 2002063030
                                               A2
                                                              20020815
                                                                                                                                     20020206
                  W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
                    RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
           US 2002128465
                                                  A1
                                                              20020912
                                                                                               US 2001-777430
                                                                                                                                     20010206
                                                                                         US 2001-777430 A 20010206
PRIORITY APPLN. INFO.:
                                                                                         US 1996-682853
                                                                                                                              A2 19960712
                                                                                         US 1999-333145
                                                                                                                              A2 19990614
```

OTHER SOURCE(S): MARPAT 137:164653 The present invention provides charge tags for attachment to materials including solid supports and nucleic acids, wherein the charge tags increase or decrease the net charge of the material. Thus, when an oligonucleotide modified with a charge tag is reduced in size (cleaved) or increased in size (elongated), the resulting product bears a net charge or a charge to mass ratio different from the original oligonucleotide thereby permitting sepn. of the original and product oligonucleotides on the basis of charge. present invention therefore further provides methods for sepg. and characterizing mols. based on the charge differentials between modified and unmodified materials, e.g., by capillary electrophoresis. Thus, MCP1 and ubiquitin transcripts were simultaneously detected in an in vitro assay using the Invader technol. and probes charge tagged with one of two phosphoramidates,

Absolute stereochemistry.
Double bond geometry unknown.

PAGE 1-A

PAGE 1-B

PAGE 2-A

H2O3PO

RN 446017-74-9 HCAPLUS

CN 3'-Uridylic acid, 5-[3-[(6-aminohexyl)amino]-3-oxo-1-propenyl]-2'-deoxy-5'-O-[[3-[2,3-dihydro-2-[3-[1-(3-hydroxypropyl)-3,3-dimethyl-3H-indolium-2-yl]-2-propenylidene]-3,3-dimethyl-1H-indol-1-yl]propoxy]hydroxyphosphinyl]uridylyl-(3'.fwdarw.5')-5-[3-[(6-aminohexyl)amino]-3-oxo-1-propenyl]-2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry unknown.

# PAGE 1-B

PAGE 2-B

L11 ANSWER 3 OF 20 HCAPLUS COPYRIGHT 2002 ACS 2002:185692 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

136:236873 Protonated antimicrobial compounds TITLE:

Dale, Roderic M. K.; Gatton, Steven L.; Arrow, INVENTOR(S):

Amy; Thompson, Terry

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 28 pp., Cont.-in-part of

U.S. Ser. No. 281,858.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO. DA	Æ
US 2002032164	A1	20020314	US 2001-847654 20	010503
US 6211349	B1	20010403	US 1998-222009 19	981230
PRIORITY APPLN. INFO.	:		US 1998-222009 A2 19	981230
			TIC 1999-281858 A2 10	990331

MARPAT 136:236873 OTHER SOURCE(S):

The present invention provides protonated compds. X-Y-Z (Y = O, P, C; X, Z = end blocking groups preventing degrdn. of the mol. and providing stability) having antimicrobial activity and a sanitizing compn. comprising a protonated compd. and a metal salt of a carboxylic acid. The protonated compds. and compns. provide efficacious antimicrobial activity against resistant strains of bacteria and opportunistic fungi. For example, the s.c. administration of compds. Nu-2, Nu-3, Nu-4, and Nu-5 (12 mg/mL) were effective in attenuating the incidence of infection of burn wounds in a mice model, a ribose deriv. Nu-4 being the most efficacious providing 100% survival. 403717-05-5 403717-06-6 403717-07-7

403717-08-8

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(protonated antimicrobial compds. and compns.)

403717-05-5 HCAPLUS RN

CN 3'-Uridylic acid, 2'-O-methyl-, mono(4-hydroxybutyl) ester, 5'-(4-hydroxybutyl hydrogen phosphate) (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 403717-06-6 HCAPLUS

CN 3'-Thymidylic acid, monobutyl ester, 5'-(butyl hydrogen phosphate) (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 403717-07-7 HCAPLUS

CN D-erythro-Pentitol, 1,4-anhydro-2-deoxy-, bis(butyl hydrogen phosphate) (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 403717-08-8 HCAPLUS

CN Phosphoric acid, P,P'-1,4-butanediyl P,P'-dibutyl ester (9CI) (CA INDEX NAME)

HCAPLUS COPYRIGHT 2002 ACS L11 ANSWER 4 OF 20 ACCESSION NUMBER: 2001:241745 HCAPLUS

134:285582 DOCUMENT NUMBER:

TITLE: Pulmonary delivery of protonated/acidified

nucleic acids

INVENTOR(S): Dale, Roderic M. K.; Gatton, Steven L.; Arrow,

PATENT ASSIGNEE(S): Oligos Etc. Inc., USA

SOURCE: U.S., 15 pp., Cont.-in-part of U.S. Ser. No.

222,009.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	PATENT NO.			KI	KIND DATE				A	PPLI	ο.	. DATE				
				B1 20010403 B1 20010403				_			_					
									WO 2000-US8244							
	W:	ΑE,	AG,	AL,	AM,	AT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,
		CR,	CU,	CZ,	DΕ,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,
		HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	ΝZ,	PL,	PT,
		RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,
		US,	UΖ,	VN,	YU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,
		DE,	DK,	ES,	FΙ,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,
		ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG	
EP 1169046				A1 20020109				E	P 20	00-9	7 20000328					
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		PT,	ΙE,	SI,	LT,	LV,	FI,	RO								
PRIORIT	Y APP	INFO	. :					US 1	998-	2220	09	A2	1998	1230		
									US 1	999-	2828	24	A	1999	0331	
									WO 2	000-	US82	44	W	2000	0328	

AΒ The present invention provides a method of treating bacterial respiratory infections by pulmonary administration of protonated/acidified nucleic acids. These modified nucleic acids are effective as bactericidal and/or bacteriostatic agents without regard to the class of bacteria, so are esp. useful when diagnosis is difficult or when multiple infectious organisms are present. The antibiotic activity of nucleic acids of the invention is not dependent on

either the specific sequence of the nucleic acid or the length of the nucleic acid mol.

331953-81-2

RL: BAC (Biological activity or effector, except adverse); BSU

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

PAGE 2-A

PAGE 2-B

REFERENCE COUNT:

24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

HCAPLUS COPYRIGHT 2002 ACS L11 ANSWER 5 OF 20

ACCESSION NUMBER: 1998:693716 HCAPLUS

DOCUMENT NUMBER: 130:52665

Optimized automated solid-phase synthesis of TITLE:

oligonucleotides and derivatives

Alvarado Urbina, Gabriel; Gruebler, Gerald; AUTHOR(S):

Weiler, Angelika; Echner, Hartmut; Stoeva, Stanka; Schernthaner, Johann; Gross, Waleri;

Voelter, Wolfgang

CORPORATE SOURCE: Biochem. Molecular Group, Eppendorf-Netheler-

Hinz G.m.b.H., Hamburg, D-22339, Germany

Zeitschrift fuer Naturforschung, B: Chemical SOURCE:

Sciences (1998), 53(9), 1051-1068

CODEN: ZNBSEN; ISSN: 0932-0776

PUBLISHER:

Verlag der Zeitschrift fuer Naturforschung

DOCUMENT TYPE: Journal English LANGUAGE:

An optimized automated synthesizer is presented for assembling oligonucleotides, thiooligonucleotides, and 5'-modified oligonucleotides including chem. phosphorylation, multihydroxyl derivatization with a non-nucleosidic phosphoramidite. The incorporation of biotin, fluorescein, and rhodamine phosphoramidites The purifn. and structure detn. of oligonucleotides is described. was confirmed using HPLC, capillary electrophoresis, and laser desorption mass spectrometry. Several applications and confirming data will be presented for gene synthesis and polymerase chain reaction expts.

ΙT 216485-54-0P 216485-55-1P

> RL: ANT (Analyte); SPN (Synthetic preparation); ANST (Analytical study); PREP (Preparation)

(optimized automated solid-phase synthesis of oligonucleotides and derivs.)

216485-54-0 HCAPLUS RN

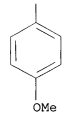
Thymidine, 5'-O-[[[5-[bis(4-methoxyphenyl)phenylmethoxy]-1-[[bis(4-CN methoxyphenyl)phenylmethoxy]methyl]pentyl]oxy]hydroxyphosphinyl]thym idylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

PAGE 2-B



RN

216485-55-1 HCAPLUS Thymidine, 5'-O-[[[5-[bis(4-methoxyphenyl)phenylmethoxy]-1-[[bis(4-methoxyphenyl)phenylmethoxy]methyl]pentyl]oxy]hydroxyphosphinyl]thymidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

308-4994 Searcher : Shears

# PAGE 1-B

IT 167212-06-8P 216485-50-6P 216485-51-7P

216485-52-8P 216485-53-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (optimized automated solid-phase synthesis of oligonucleotides and derivs.)

RN 167212-06-8 HCAPLUS

CN Thymidine, 5'-O-[hydroxy[[5-hydroxy-1-(hydroxymethyl)pentyl]oxy]phos phinyl]thymidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 216485-50-6 HCAPLUS

CN Thymidine, 2'-deoxy-5'-O-[hydroxy[[5-hydroxy-1-(hydroxymethyl)pentyl]oxy]phosphinyl]adenylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 216485-51-7 HCAPLUS

CN Cytidine, 2'-deoxy-5'-O-[hydroxy[[5-hydroxy-1-(hydroxymethyl)pentyl]oxy]phosphinyl]cytidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxy-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 216485-52-8 HCAPLUS

CN Thymidine, 5'-O-[hydroxy[[5-hydroxy-1-(hydroxymethyl)pentyl]oxy]phos phinyl]thymidylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 216485-53-9 HCAPLUS
CN Thymidine, 5'-O-[hydroxy[[5-hydroxy-1-(hydroxymethyl)pentyl]oxy]phos phinyl]thymidylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 6 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: DOCUMENT NUMBER:

1997:26983 HCAPLUS

126:114175

TITLE:

Targeted cleavage of RNA using antisense oligonucleotide linked to 2',5'-oligoadenylate

activator of RNase

INVENTOR(S):

Torrence, Paul; Silverman, Robert; Maitra,

Ratan; Lesiak, Krystyna

PATENT ASSIGNEE(S):

Cleveland Clinic Foundation and National

Institutes of Health, USA

SOURCE:

U.S., 43 pp., Cont.-in-part of U.S. Ser. No.

965,666, abandoned.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PA'	TENT :	NO.		KI	ΝD	DATE			A	PPLI	CATI	ON NO	ο.	DATE		
US CA WO	CA 2147282			A0 19930401 AA 19940428				US 1992-965666						1993 1992 1993 1993		
		AT,	CA, BE,		DE,	DK,	ES,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,
	9455 6692			A: B:		1994 1996			А	บ 19	94-5	5858		1993	1020	
	6669 6669	10		B	1	1995 2002	0130							1993		
7.0		PT,	SE											LU,		NL,
AT	0850: 2126 5677:	64				1996 2002 1997	0215		A	Р 19 Т 19 Ѕ 19	94-9	01178	3	1993 1993 1995	1020	
	6271	369		B	1	2001	0807	1	U US 1	S 19 992-	97-9! 9656	5019 66	6 B2	1997 1992	1014	
								1	WO 1	993- 993- 995-	US10:	103	M	1993 1993 1995	1020	

AΒ A method of using a chimeric mol. made up of an antisense oligonucleotide attached to a 2',5'-oligoadenylate mol. to specifically cleave a sense strand of RNA, wherein the antisense oligonucleotide of the chimeric mol. is hybridized to the sense strand of RNA in the presence of 2',5'-dependent RNase is claimed. Chimeric 2',5'-oligoA-antisense oligonucleotides were synthesized and tested in vitro and in vivo. The chimeric mol. caused sequence-specific cleavage. The 2',5'-oligoA moiety enhanced the ability of antisense oligonucleotides to inhibit specific gene expression. When administered to mammalian cells, destruction of specific RNA mols. occurred without having to treat the cells in any special way in order to get the chimeric mols. into the cell.

TΤ 185997-79-9P

> RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(targeted cleavage of RNA using antisense oligonucleotide linked to 2',5'-oligoadenylate activator of RNase) 185997-79-9 HCAPLUS

RN

CN

Thymidine, 5'-O-phosphonoadenylyl-(2'.fwdarw.5')-adenylyl-(2'.fwdarw.5')-adenylyl-(2'.fwdarw.5')-adenylyloxyphosphinicooxy-1,4butanediyloxyphosphinicooxy-1,4-butanediyloxyphosphinico-(2'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

## Absolute stereochemistry.

PAGE 1-A H2O3PO R R НО NH2 NH<sub>2</sub>

## PAGE 1-B

Shears 308-4994 Searcher

PAGE 2-C

HCAPLUS COPYRIGHT 2002 ACS L11 ANSWER 7 OF 20

ACCESSION NUMBER:

1996:487020 HCAPLUS

DOCUMENT NUMBER:

125:214989

TITLE:

AUTHOR(S):

Conjugates of minor groove DNA binders with oligodeoxynucleotides: synthesis and properties

Levina, Asya S.; Metelev, Valeri G.; Cohen, Aharon S.; Zamecnik, Paul C.

CORPORATE SOURCE:

Worcester Foundation Biomed. Res., Shrewsbury,

MA, 01545-2737, USA

SOURCE: Antisense & Nucleic Acid Drug Development

(1996), 6(2), 75-85

CODEN: ANADF5; ISSN: 1087-2906

PUBLISHER: Liebert
DOCUMENT TYPE: Journal
LANGUAGE: English

Oligodeoxynucleotide conjugates of netropsin (Nt) and distamycin A AR (Dst) were synthesized, and the thermal stability of several model DNA duplexes contg. conjugates was studied. Two Dst residues conjugated at both ends of the oligodeoxynucleotide were needed for substantial increase in the melting temp. of the corresponding duplex (.DELTA.Tm > 30.degree.). Two attached Dst residues had a greater effect on the Tm value than did two free mols. of Dst per duplex. In contrast to Dst, one Nt mol. linked to the oligonucleotide was enough to influence the thermal stability of the duplexes. Like Dst, the attached Nt appeared to stabilize duplexes much more than free Nt mols. Attachment of Nt to either the 5'- or 3'-end of the different nonadeoxynucleotides contg. 5'...TTAAA... or 5'... TATA sites increased Tm of their duplexes by 21.degree.-25.degree., whereas .DELTA.Tm for free Nt was 8.degree.-15.degree. (.DELTA..DELTA.Tm = 10.degree.-14.degree.). The same phenomenon was shown for oligonucleotide phosphorothioates (.DELTA.Tm were 18.degree.-22.degree. and 9.degree.-13.degree. for attached and free Nt, resp.; .DELTA..DELTA.Tm = 9.degree.). This effect was even more pronounced for a hairpin oligonucleotide (.DELTA..DELTA.Tm = 18.degree.).

### IT 180840-08-8

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(synthesis and properties of conjugates of minor groove DNA binders distamycin and netropsin with oligodeoxynucleotides)

RN 180840-08-8 HCAPLUS

CN 3'-Cytidylic acid, 2'-deoxy-5'-O-(1,21,21-trihydroxy-1,21-dioxido-2,5,8,11,14,17,20-heptaoxa-1,21-dipnosphaheneicos-1-yl)cytidylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxy-, 3'-(19,19-dihydroxy-19-oxido-3,6,9,12,15,18-hexaoxa-19-phosphanonadec-1-yl) ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

# PAGE 1-A

# PAGE 1-B

Searcher : Shears

.308-4994

PAGE 2-D

L11 ANSWER 8 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1996:286949 HCAPLUS

DOCUMENT NUMBER: 124:334809

TITLE: Stable triple helixes formed by oligonucleotide

N3' .fwdarw. P5' phosphoramidates inhibit

transcription elongation

AUTHOR(S): Escude, Christophe; Giovannangeli, Carine; Sun,

Jian-Sheng; Lloyd, David H.; Chen, Jer-Kang;
Gryaznov, Sergei M.; Garestier, Therese; Helene,

Claude

CORPORATE SOURCE: Lab. Biophys., Cent. Natl. Rech. Sci., Paris,

75231, Fr.

SOURCE: Proceedings of the National Academy of Sciences

of the United States of America (1996), 93(9),

4365-4369

CODEN: PNASA6; ISSN: 0027-8424

PUBLISHER: National Academy of Sciences

DOCUMENT TYPE: Journal LANGUAGE: English

Oligonucleotide analogs with N3' .fwdarw. P5' phosphoramidate linkages bind to the major groove of double-helical DNA at specific oligopurine-oligopyrimidine sequences. These triple-helical complexes are much more stable than those formed by oligonucleotides with natural phosphodiester linkages. Oligonucleotide phosphoramidates contg. thymine and cytosine or thymine, cytosine, and guanine bind strongly to the polypurine tract of human immunodeficiency virus proviral DNA under physiol. conditions. Site-specific cleavage by the Dra I restriction enzyme at the 5' end of the polypurine sequence was inhibited by triplex formation. A eukaryotic transcription assay was used to investigate the effect of oligophosphoramidate binding to the polypurine tract sequence on transcription of the type 1 human immunodeficiency virus nef gene under the control of a cytomegalovirus promoter. An efficient arrest of RNA polymerase II was obsd. at the specific triplex site at submicromolar concns.

### IT 119874-42-9

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological

```
study)
          (Acr-d(T-T-T-C-C-T-C-T-C-T); stable triple helixes formed by
          oligonucleotide N3' .fwdarw. P5' phosphoramidates inhibit
          transcription elongation)
RN 119874-42-9 HCAPLUS
CN 5'-Thymidylic acid, thymidylyl-(5'.fwdarw.3')-2'-deoxycytidylyl-
          (5'.fwdarw.3')-thymidylyl-(5'.fwdarw.3')-2'-deoxycytidylyl-
          (5'.fwdarw.3')-2'-deoxycytidylyl-(5'.fwdarw.3')-thymidylyl-
          (5'.fwdarw.3')-2'-deoxycytidylyl-(5'.fwdarw.3')-2'-deoxycytidylyl-
          (5'.fwdarw.3')-thymidylyl-(5'.fwdarw.3')-thymidylyl-(5'.fwdarw.3')-,
          5'-[5-[(6-chloro-2-methoxy-9-acridinyl)amino]pentyl] ester (9CI)
          (CA INDEX NAME)
```

PAGE 1-A

PAGE 2-B

PAGE 3-A

L11 ANSWER 9 OF 20 HCAPLUS COPYRIGHT 2002 ACS 1995:767385 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

TITLE:

123:286546

Synthesis of nucleic acid probes for sensing a

DNA and RNA molecules

INVENTOR(S):

De Vos, Marie-Joelle; Bollen, Alex

PATENT ASSIGNEE(S):

La Region Wallonne, Belg.

SOURCE:

PCT Int. Appl., 97 pp.

CODEN: PIXXD2

Searcher :

Shears 308-4994 DOCUMENT TYPE:

Patent English

LANGUAGE:

GI

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.			KIND DATE				APPLICATION NO.							DATE				
		119364 119364							WO	19	94-B	E13		1994	0218	3		
							ES,	FR,	GE	3,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	
BE	10079	_		A	3	1995	51121			BE	19	93-1	60		1993	0219		
CA	21557	57		$\mathbf{A}$	A	1994	10901			CA	. 19	94-23	1557	57	1994	0218		
EP	68495	4		A	1	1995	1206			ΕP	19	94-90	0609	9	1994	0218		
EP	68495	4		B	1	1998	30722											
	R:	ΑT,	BE,	CH,	DE,	ES,	FR,	GB,	ΙT	٠,	LI,	NL						
JP	08511	678		T	2	1996	51210			JР	19	94-5	1849	2	1994	0218		
AT	16869	6		Ε		1998	30815			ΑT	19	94-90	0609	9	1994	0218		
ES	21222	33		T	3	1998	31216			ES	19	94-90	0609	9	1994	0218		
NO	95032	51		Α		1995	0818			NO	19:	95-32	251		1995	0818		
FI	95038	97		Α		1995	1002			FI	19	95-38	397		1995	0818		
US	59691	28		Α		1999	1019			US	19	95-50	728	3	1995	0821		
PRIORITY	Y APPL	Ν. :	INFO	. :					ΒE	19	93-	160			1993	0219		
									WO	19	94-1	BE13			1994	0218		
OTHER SO	OURCE (	S):			MAR	PAT	123:2	2865	46									

 $[(M)Y^{1}-(L)X^{1}]Z^{1}-S[(L)X-(M)Y]Z$ 

Nucleic acid probes for sensing a DNA or RNA mol. The probe I AB comprises (a) an oligodeoxyribonucleotide or oligodeoxyribonucleotide portion consisting of a DNA or RNA nucleic acid sequence (S) depending of the kind of mol. to be sensed, and (b) a non-nucleotide portion having chem. properties enabling the direct or indirect attachment of one or more detection units or biotin labeling elements (M) that are non-isotopically detectable by oligodeoxyribonucleotide. of color or light, and chem. arm (L), with x, x1, z, z1 are > or equal to zero, y and y1 are never both equal to zero. The probe is characterized in that portion (b) consists of a chain of phosphate units between which are inserted alkyl units, i.e. (bl) a no. of alkyl units joining the various phosphate groups and having no particular functionality, and (b2) alkyl units having primary amine functions enabling coupling to various reagents and thus direct or indirect detection, units (b2) being bound to portion (a) or sequence (S) via units (b1). Sequence (S) is linked at its 5' and/or 3' end ot one or more labeling elements (M). Such probes are useful for sensing and diagnosing hereditary genetic diseases, oncogenes, and viral, bacterial or parasitic diseases.

IT 166887-51-0P 166887-52-1P 166887-53-2P 167211-99-6P 167212-03-5P 167212-04-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. of biotin labeled nucleic acid probes for sensing a DNA

and RNA mols.)

RN 166887-51-0 HCAPLUS

CN Thymidine, 5'-O-[hydroxy(3-hydroxy-1-methylpropoxy)phosphinyl]thymid yly1-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 166887-52-1 HCAPLUS

CN Thymidine, 5'-O-[[1-[[(6-amino-1-oxohexyl)amino]methyl]-3-hydroxypropoxy]hydroxyphosphinyl]thymidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

RN 166887-53-2 HCAPLUS
CN Thymidine, 5'-O-[[[15-amino-5-hydroxy-7-(2-hydroxyethyl)-1-methyl-5-oxido-10-oxo-4,6-dioxa-9-aza-5-phosphapentadec-1-yl]oxy]hydroxyphosphinyl]thymidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

RN 167211-99-6 HCAPLUS

CN Thymidine, 5'-O-[[[7-[[(6-amino-1-oxohexyl)amino]methyl]-5,11,15-trihydroxy-1,13-dimethyl-5,11-dioxido-4,6,10,12-tetraoxa-5,11-diphosphapentadec-1-yl]oxy]hydroxyphosphinyl]thymidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

RN 167212-03-5 HCAPLUS

CN Thymidine, 5'-O-[hydroxy[3-[[hydroxy(3-hydroxy-1-methylpropoxy]phosphinyl]oxy]-1-methylpropoxy]phosphinyl]thymidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

RN 167212-04-6 HCAPLUS

CN 3'-Thymidylic acid, 5'-O-[[3-[[[1-[[(6-amino-1-oxohexyl)amino]methyl]-3-hydroxypropoxy]hydroxyphosphinyl]oxy]-1-methylpropoxy]hydroxyphosphinyl]thymidylyl-(3'.fwdarw.5')-, 3'-[3-[[[4-[(6-amino-1-oxohexyl)amino]-3-[[hydroxy(3-hydroxybutoxy)phosphinyl]oxy]butoxy]hydroxyphosphinyl]oxy]butyl]ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

## IT 166887-47-4P 167212-06-8P 167212-07-9P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (prepn. of biotin labeled nucleic acid probes for sensing a DNA
 and RNA mols.)

RN 166887-47-4 HCAPLUS

CN Thymidine, 5'-O-[[[5-[[[(6-aminohexyl)oxy]hydroxyphosphinyl]oxy]-1[[[[(6-aminohexyl)oxy]hydroxyphosphinyl]oxy]methyl]pentyl]oxy]hydrox
yphosphinyl]thymidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

RN 167212-06-8 HCAPLUS

CN Thymidine, 5'-O-[hydroxy[[5-hydroxy-1-(hydroxymethyl)pentyl]oxy]phos phinyl]thymidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

$$\begin{array}{c} \text{OH} \\ \text{HO} \\ \text{O} \\$$

RN 167212-07-9 HCAPLUS

Thymidine, 5'-O-[hydroxy[[5-[[hydroxy[[5-hydroxy-1-(hydroxymethyl)pentyl]oxy]phosphinyl]oxy]-1-[[[hydroxy[[5-hydroxy-1-(hydroxymethyl)pentyl]oxy]phosphinyl]oxy]methyl]pentyl]oxy]phosphinyl]thymidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

#### PAGE 1-A

PAGE 1-B

L11 ANSWER 10 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1995:46745 HCAPLUS

DOCUMENT NUMBER:

122:127234

TITLE:

A novel fluorogenic substrate for ribonucleases.

Synthesis and enzymic characterization

AUTHOR(S):

Zelenko, Ottilie; Neumann, Ulf; Brill, Wolfgang;

Pieles, Uwe; Moser, Heinz E.; Hofsteenge, Jan

CORPORATE SOURCE:

Friedrich Miescher-Institut, Basel, CH-4002,

SOURCE:

Nucleic Acids Research (1994), 22(14), 2731-9

CODEN: NARHAD; ISSN: 0305-1048

DOCUMENT TYPE:

Journal

LANGUAGE: English The synthesis and enzymic characterization of 5'-O-[4-(2,4dinitrophenylamino)butyl]phosphoryluridylyl-(3'.fwdarw.5')-2'deoxyadenosine-3'-[N-[(2-aminobenzoyl)aminoprop-3-yl]phosphate] (DUPAAA), a novel fluorogenic substrate for pancreatic-type RNases is described. It consists of the dinucleotide, uridylyl-3',5'deoxyadenosine, to which a fluorophore, o-aminobenzoic acid, and a quencher, 2,4-dinitroaniline, have been attached by means of phosphodiester linkages. Due to intramol. quenching, the intact substrate displayed very little fluorescence. Cleavage of the phosphodiester bond at the 3'-side of the uridylyl residue by RNase caused a 60-fold increase in fluorescence. This allowed the continuous and highly sensitive monitoring of enzyme activity. The substrate was turned over efficiently by RNases of the pancreatic type, but no cleavage was obsd. with microbial RNase T1. Compared to the dinucleotide substrate, UpA, the specificity const. with RNase A, RNase PL3, and RNase Us increased 6-, 18-, and 29-fold, resp. These differences in increased catalytic efficiency most likely reflect differences in the importance of subsites on the enzyme in the binding of elongated substrates. Studies on the interactions of RNase inhibitor with RNase A using DUPAAA as a reporter substrate showed that it was well suited for monitoring this very tight protein-protein interaction using pre-steady-state kinetic methods.

#### 161054-62-2P TT

RL: BPR (Biological process); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); PROC (Process)

(synthesis and enzymic characterization of novel fluorogenic substrate for RNases)  $\,$ 

RN 161054-62-2 HCAPLUS

CN 3'-Adenylic acid, 5'-O-[[4-[(2,4-dinitrophenyl)amino]butoxy]hydroxyp hosphinyl]uridylyl-(3'.fwdarw.5')-2'-deoxy-, 3'-[3-[(2-aminobenzoyl)amino]propyl] ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

L11 ANSWER 11 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1994:98485 HCAPLUS

DOCUMENT NUMBER:

120:98485

TITLE:

Improvements in or relating to DNA cloning techniques and products for use therewith

#### 09/847654

INVENTOR(S): Taylor, Philip Neil PATENT ASSIGNEE(S): University of Hull, UK SOURCE: PCT Int. Appl., 46 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9319186	A1	19930930	WO 1993-GB584	19930322
•		, DK, ES, F	CR, GB, GR, IE, IT, LU	, MC, NL, PT,
SE EP 631625	A1	19950104	EP 1993-906721	19930322
R: CH, DE US 5604122	, FR, GB A	•	US 1994-307713	19941114
PRIORITY APPLN. INF	·O.:		GB 1992-6210 WO 1993-GB584	19920321 19930322

AB A method of cloning a foreign DNA into a DNA vector is described. The method comprises ligating (1) a DNA vector having a single-standed (ss) DNA overhang at each end, said overhangs being mutually incompatible so as to prevent self-religation; with (2) a linear piece of foreign DNA having a ss-DNA overhang at each end, each foreign DNA overhang being complementary to but at least 1 base shorter than each of the vector overhangs and being capable of base-pairing along the entire length of the overhang with 1 of the vector overhangs; followed by sealing the gap by either transforming the double-stranded DNA having a gap therein into a bacterium or transfering it into a bacterium after packaging into a bacteriophage. A DNA cloning kit comprising a defined DNA vector, DNA linkers, a restriction endonuclease, and a DNA ligase is also claimed.

IT 151837-15-9

RL: BIOL (Biological study)

(linker, cloning vector contg.)

RN 151837-15-9 HCAPLUS

CN Guanosine, 2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxy-3'-de(phosphinicooxy)-5-methylcytidylylmethyleneoxyphosphinico-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')-2'-deoxy-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

# PAGE 1-B

Searcher : She

Shears 308-4994

PAGE 2-B

L11 ANSWER 12 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1991:58069 HCAPLUS

DOCUMENT NUMBER: 114:58069

TITLE: Highly selective affinity labeling of an E. coli

RNA polymerase promoter complex with reactive

derivatives of oligonucleotide primers of

various chemical specificities

AUTHOR(S): Tsarev, I. G.; Mustaev, A. A.; Zaychikov, E. F.;

Alikina, T. Yu.; Ven'yaminova, A. G.; Repkova,

M. N.

CORPORATE SOURCE: Limnol. Inst., Irkutsk, USSR

SOURCE: Bioorg. Khim. (1990), 16(6), 765-79

CODEN: BIKHD7; ISSN: 0132-3423

DOCUMENT TYPE: Journal LANGUAGE: Russian

AB A technique of highly selective affinity labeling, which includes covalent modification of the enzyme-T7A2 promoter complex with reactive oligonucleotide derivs. and subsequent elongation of the attached oligonucleotide residue with a radioactive substrate, was used to study the product-binding site of Escherichia coli RNA polymerase. Different oligonucleotides complementary to the T7A2 promoter (with lengths of 2-8 residues) contg. 5'-terminal phosphorylating, alkylating, or aldehyde groups were used for labeling. The procedure resulted in labeling DNA and .beta., .beta.', or .sigma. subunits of the enzyme, which are therefore believed to contact growing RNA in the course of initiation. Consideration of the labeling patterns as a function of oligonucleotide length as well as of the structure and chem. specificity of the reactive groups led to a tentative topog. scheme of the RNA polymerase product-binding region.

IT 131401-19-9P 131401-20-2P 131401-22-4P 131401-23-5P 131401-24-6P 131401-25-7P 131419-77-7P 131419-78-8P 131419-79-9P

131419-80-2P

RL: SPN (Synthetic preparation); PREP (Preparation)

## 09/847654

PAGE 1-A

0

PAGE 3-A

131401-20-2 HCAPLUS RN

Cytidine, 2'-deoxy-5'-O-[[2-[(4-formylphenyl)methylamino]ethoxy]hydr oxyphosphinyl]adenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')-(9CI) (CA INDEX NAME) CN

o=

PAGE 2-B

PAGE 3-A | | |NH2

PAGE 4-A

RN 131401-22-4 HCAPLUS
CN Cytidine, 5'-O-[[2-[(2-chloroethyl)(4-formylphenyl)amino]ethoxy]hydr
oxyphosphinyl]thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX
NAME)

PAGE 1-A

$$\begin{array}{c} & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

СНО

PAGE 2-B

RN 131401-23-5 HCAPLUS
CN Cytidine, 5'-O-[[2-[(2-chloroethyl)(4-formylphenyl)amino]ethoxy]hydr
oxyphosphinyl]-2'-deoxyadenylyl-(3'.fwdarw.5')-thymidylyl(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-

# (3'.fwdarw.5') - (9CI) (CA INDEX NAME)

## PAGE 1-A

OHC
$$N - CH_2 - CH_2 - O - P - O - CH_2$$

$$C1CH_2 - CH_2$$

$$HO - P = O$$

# PAGE 2-A

Searcher : Shears

308-4994

PAGE 2-B

PAGE 3-B

RN 131401-24-6 HCAPLUS
CN Cytidine, 5'-O-[[2-[(2-chloroethyl)(4-formylphenyl)amino]ethoxy]hydr
oxyphosphinyl]-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX
NAME)

# PAGE 1-A

# PAGE 1-B

PAGE 2-B

PAGE 3-A

O
N
Me
O

RN 131401-25-7 HCAPLUS
CN Cytidine, 5'-O-[[2-[(2-chloroethyl)(4-formylphenyl)amino]ethoxy]hydr
oxyphosphinyl]-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-thymidylyl(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')-2'-deoxycytidylyl(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

PAGE 1-B

0=

Me 
$$\stackrel{\text{H}}{\longrightarrow} \stackrel{\text{O}}{\longrightarrow} \stackrel{\text{OH}}{\longrightarrow} \stackrel{\text$$

PAGE 2-B

Searcher :

Shears

308-4994

PAGE 3-A NH2

NH<sub>2</sub>

PAGE 4-A

RN

131419-77-7 HCAPLUS
Cytidine, 5'-O-[[2-[(4-formylphenyl)methylamino]ethoxy]hydroxyphosph
inyl]thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME) CN

PAGE 1-A

$$\begin{array}{c} & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

PAGE 1-B

308-4994

PAGE 2-B

СНО

RN 131419-78-8 HCAPLUS
CN Cytidine, 5'-O-[[2-[(4-formylphenyl)methylamino]ethoxy]hydroxyphosph
inyl]-2'-deoxyadenylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI)
(CA INDEX NAME)

PAGE 1-A

OHC
$$N - CH_2 - CH_2 - O - P - O - CH_2$$

$$HO - P = O$$

CH<sub>2</sub>

# PAGE 2-B

PAGE 3-B

RN 131419-79-9 HCAPLUS
CN Cytidine, 5'-O-[[2-[(4-formylphenyl)methylamino]ethoxy]hydroxyphosph
inyl]-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX
NAME)

PAGE 1-A

PAGE 2-B

RN 131419-80-2 HCAPLUS
CN Cytidine, 5'-O-[[2-[(2-chloroethyl)(4-formylphenyl)amino]ethoxy]hydr
oxyphosphinyl]-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

PAGE 3-A

HCAPLUS COPYRIGHT 2002 ACS L11 ANSWER 13 OF 20

ACCESSION NUMBER: 1991:58051 HCAPLUS

DOCUMENT NUMBER: 114:58051

TITLE: Stereochemical studies of the .beta.-elimination

> reactions at aldehydic abasic sites in DNA: endonuclease III from Escherichia coli, sodium

hydroxide, and Lys-Trp-Lys

Mazumder, Abhijit; Gerlt, John A.; Absalon, AUTHOR(S):

Michael J.; Stubbe, JoAnne; Cunningham, Richard

P.; Withka, Jane; Bolton, Philip H.

Dep. Chem., Massachusetts Inst. Technol., CORPORATE SOURCE:

Cambridge, MA, 02139, USA Biochemistry (1991), 30(4), 1119-26 SOURCE:

CODEN: BICHAW; ISSN: 0006-2960

DOCUMENT TYPE: Journal LANGUAGE: English

The DNA strand cleavage reaction catalyzed by endonuclease III from E. coli (endo III) on the 3'-side of aldehydic abasic sites proceeds by a syn .beta.-elimination involving abstraction of the 2'-pro-S proton and formation of a trans .alpha.,.beta.-unsatd. aldose product; the same stereochem. course was previously reported for the reaction catalyzed by UV endonuclease V from bacteriophage T4 (UV endo V). Since UV endo V does not contain a 4Fe-4S center, the 4Fe-4S center present in endo III need not be assigned a unique role in the .beta.-elimination reaction. The .beta.-elimination reactions that occur under alk. conditions (0.1 N NaOH) and in the presence of the tripeptide Lys-Trp-Lys proceed by anti .beta.-elimination mechanisms involving abstraction of the 2'-pro-R proton and formation of a trans .alpha.,.beta.-unsatd. aldose product. The different stereochem. outcomes of the enzymic and nonenzymic .beta.-elimination reactions support the hypothesis that the enzyme-catalyzed reactions may involve general-base-catalyzed abstraction of the 2'-pro-S proton by the internucleotidic phosphodiester leaving group.

IT130882-87-0

RL: RCT (Reactant)

(reaction of, with endonuclease III of Escherichia coli, stereochem. and mechanism of, nonenzymic reaction in relation to)

RN130882-87-0 HCAPLUS

CN DNA, d(C-G-C-A-G-(3'.fwdarw.5)-(oxyphosphinico-2-deoxy-D-erythropentos-3-O-ylphosphinicooxy)-(3.fwdarw.5')-C-A-G-C-C), complex with DNA d(G-G-C-T-G-A-C-T-G-C-G) (1:1) (9CI) (CA INDEX NAME)

CM1

CRN 130882-86-9 CMF C107 H135 N43 O65 P10

Absolute stereochemistry.

PAGE 3-B

PAGE 4-A

CM

CRN

130882-85-8 C100 H130 N40 O62 P10 CMF

PAGE 1-B

Searcher : Shears

308-4994

$$H_2N$$
 $N$ 
 $O$ 
 $CH_2$ 
 $O$ 
 $P$ 
 $O$ 
 $O$ 

Searcher

Shears

08-4994

PAGE 3-A

#### 09/847654

PAGE 4-A

L11 ANSWER 14 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1989:439835 HCAPLUS

DOCUMENT NUMBER:

111:39835

TITLE:

Preparation of .alpha.-D-oligonucleotide

derivatives as artificial nucleases

INVENTOR(S):

Helene, Claude; Nguyen, Thank Thuong Centre National de la Recherche Scientifique,

PATENT ASSIGNEE(S):

Fr.

SOURCE:

Fr. Demande, 35 pp.

CODEN: FRXXBL

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
	A1		FR 1986-16797	19861202	
	B1 A1	19900413 19880616	WO 1987-FR481	19871202	
W: JP, US	CH. DE	, FR, GB, IT,	LII. NI. SE		
EP 290583	A1	19881117	EP 1988-900022	19871202	
		19930407 , FR, GB, IT,	LI, LU, NL, SE		
			JP 1988-500386 AT 1988-900022	19871202 19871202	
PRIORITY APPLN. INFO			FR 1986-16797	19861202	
			FR 1987-4339 EP 1988-900022	19870327 19871202	
WO 1987-FR481 19871202 OTHER SOURCE(S): MARPAT 111:39835					

GΙ

AΒ The title compds. [I;  $B = nucleic \ acid \ base; \ X = O-, S-, \ alkyl,$ alkoxy, etc.; L = O, S, NH; J = H, OH; n = integer, O; R, R1 = H, YZ, Y1Z1; Y, Y1 = alkylene (alk), P(O)XS-, P(O)X-O-alk-, etc.; Z, Z1 = intercalating agent, a group that may directly or indirectly link with nucleotides], useful as artificial nucleases (no data), are prepd. Octanucleotide .alpha.-(Tp)8(CH2)5-Acr [Acr = (2-chloro-6-methoxy-10-acridinyl)amino] was prepd. in many steps from 5'-O-(dimethoxytrityl)-.alpha.-thymidine. IT112591-87-4P 112591-90-9P 119051-51-3P 119082-35-8P 120886-02-4P RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, as artificial nuclease) RN 112591-87-4 HCAPLUS .alpha.-Thymidine, 5'-O-[[[5-[(6-chloro-2-methoxy-9-CN acridinyl)amino]pentyl]oxy]hydroxyphosphinyl]-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-(9CI) (CA INDEX NAME)

Me 
$$\stackrel{\text{H}}{\longrightarrow}$$
  $\stackrel{\text{O}}{\longrightarrow}$   $\stackrel{\text{CH}_2-\text{O}-\text{P}}{\longrightarrow}$   $\stackrel{\text{O}}{\longrightarrow}$   $\stackrel{\text{CH}_2-\text{O}-\text{P}}{\longrightarrow}$   $\stackrel{\text{O}}{\longrightarrow}$   $\stackrel{\text{CH}_2-\text{O}}{\longrightarrow}$   $\stackrel{\text{C$ 

PAGE 1-B

Searcher :

Shears

308-4994

PAGE 2-A

Cl

PAGE 2-B

PAGE 3-A

Me 
$$CH_2-O-P-O-O$$
 $Me$ 
 $N+O$ 
 $O+O$ 
 $O+O$ 

#### 09/847654

RN 112591-90-9 HCAPLUS

CN .alpha.-Thymidine, 5'-O-[[[5-[(6-chloro-2-methoxy-9-acridinyl)amino]pentyl]oxy]hydroxyphosphinyl]-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-, 3'-[O-[2-(4-azidophenyl)-2-oxoethyl] hydrogen phosphorothioate] (9CI) (CA INDEX NAME)

PAGE 1-A

Me

O

$$\begin{array}{c}
H \\
N \\
O
\end{array}$$
 $\begin{array}{c}
CH_2 - O \\
O \\
OH
\end{array}$ 
 $\begin{array}{c}
CH_2 - O \\
OH
\end{array}$ 
 $\begin{array}{c}
CH_2 - O \\
OH
\end{array}$ 
 $\begin{array}{c}
CH_2 - O \\
OH
\end{array}$ 

PAGE 2-A

PAGE 2-B

PAGE 3-A

RN 119051-51-3 HCAPLUS

.alpha.-Thymidine, 5'-O-[[[5-[(6-chloro-2-methoxy-9-acridinyl)amino]pentyl]oxy]hydroxyphosphinyl]-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-, 3'-(dihydrogen phosphorothioate) (9CI) (CA INDEX NAME)

## PAGE 1-A

## PAGE 1-B

Searcher :

Shears

308-4994

Cl

$$\begin{array}{c} & & & & & \\ & & & & \\ & & & & \\$$

# PAGE 2-B

## PAGE 3-A

RN 119082-35-8 HCAPLUS

.alpha.-5'-Cytidylic acid, .alpha.-thymidylyl-(5'.fwdarw.3')-.alpha.thymidylyl-(5'.fwdarw.3')-2'-deoxy-.alpha.-adenylyl-(5'.fwdarw.3').alpha.-thymidylyl-(5'.fwdarw.3')-2'-deoxy-.alpha.-adenylyl(5'.fwdarw.3')-.alpha.-thymidylyl-(5'.fwdarw.3')-.alpha.-thymidylyl(5'.fwdarw.3')-2'-deoxycytidylyl-(5'.fwdarw.3')-2'-deoxy-,
5'-[5-[(6-chloro-2-methoxy-9-acridinyl)amino]pentyl] ester (9CI)
(CA INDEX NAME)

#### PAGE 1-A

PAGE 2-B

\_\_\_\_OMe

PAGE 3-B

PAGE 4-A

O
N
HN
Me

```
RN 120886-02-4 HCAPLUS
CN .alpha.-Thymidine, 5'-O-[[[5-[(6-chloro-2-methoxy-9-acridinyl)amino]pentyl]oxy]hydroxyphosphinyl]-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-
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(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-, 3'-[O-(carboxymethyl) hydrogen phosphorothioate] (9CI) (CA INDEX NAME)

PAGE 1-A

## PAGE 3-A

PAGE 4-A

L11 ANSWER 15 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1989:135661 HCAPLUS

DOCUMENT NUMBER: 110:135661

TITLE: Preparation of .alpha.-(deoxy)oligonucleotide

derivatives conaining an intercalating agent or

reactive radical

INVENTOR(S): Helene, Claude; Imbach, Jean Louis; Nguyen Thanh

Thuong; Paoletti, Claude; Rayner, Bernard

PATENT ASSIGNEE(S): Centre National de la Recherche Scientifique,

Fr.

SOURCE: PCT Int. Appl., 111 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent French

LANGUAGE: French

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	A1	19880616	WO 1987-FR481	19871202
W: JP, US RW: AT, BE,	CH, DE	, FR, GB,	IT, LU, NL, SE	
FR 2607507	A1	19880603	FR 1986-16797	19861202
FR 2607507	B1	19900413		
FR 2612930	A1	19880930	FR 1987-4339	19870327
FR 2612930	В1	19901228		
JP 01502187	T2	19890803	JP 1988-500386	19871202
AT 87932	E	19930415	AT 1988-900022	19871202
PRIORITY APPLN. INFO	. :		FR 1986-16797	19861202
			FR 1987-4339	19870327
			EP 1988-900022	19871202
			WO 1987-FR481	19871202
OTHER SOURCE(S):	CA	SREACT 110	:135661; MARPAT 110:13	5661

II

AΒ The title compds. [I; B = optionally substituted nucleotide base; J = H, OH; R, R1 = H, YZ, Y1Z1; n = 0, an integer; L = O, S, NH; Y, Y1 = alkylene, carbonylalkylene, carbonyliminoalkylene, etc.; Z, Z1 = intercalating agent residue, radical carrier of a reactive function], useful as antibiotics, antiviral, antitumor, or antiparacitic agents, are prepd. .alpha.-[D(CATGCG)] was prepd. by coupling of the appropriate 5'-O-dimethoxytrityl-.alpha.-nucleoside 3'-O-[(2-chloro-4-tritylphenyl)(2-cyanoethyl)] phosphates (obtained via, e.g., transglycosylation-anomerization of 4-N-benzoyl-3',5'-di-O-acetyl-2'-deoxycytidine) via the conventional detritylation, decyanoethylation, and condensation steps in as many cycles as necessary. Dmtr-.alpha.(TTTT) (Dmtr = dimethoxytrityl) was condensed with Dmtr-NH(CH2)5CO2H in the presence of DCC and DMAP to give Dmtr-.alpha.(TTTT)-02C(CH2)5NH-Dmtr, which was deprotected with 80% AcOH to give .alpha.(TTTT)-O2C(CH2)5NH2, which was condensed with an enzyme-oxidized deriv. of 2-methyl-9-hydroxyellipticinium acetate to give a fluorescent compd. (II).

112591-87-4P 119051-51-3P 119051-52-4P TΤ 119051-53-5P 119082-35-8P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (prepn. of, as antitumor, antimicrobial, and antiparasitic agent)

RN

112591-87-4 HCAPLUS .alpha.-Thymidine, 5'-O-[[[5-[(6-chloro-2-methoxy-9-CN acridinyl)amino]pentyl]oxy]hydroxyphosphinyl]-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5') - (9CI) (CA INDEX NAME)

## PAGE 1-A

$$\begin{array}{c} & & & \\ & &$$

# PAGE 1-B

PAGE 2-A

Cl

PAGE 2-B

PAGE 3-A

Me 
$$\stackrel{\text{H}}{\longrightarrow}$$
  $\stackrel{\text{O}}{\longrightarrow}$   $\stackrel{\text{OH}}{\longrightarrow}$   $\stackrel{\text{CH}_2}{\longrightarrow}$   $\stackrel{\text{CH}_2}{\longrightarrow}$   $\stackrel{\text{O}}{\longrightarrow}$   $\stackrel{\text{CH}_2}{\longrightarrow}$   $\stackrel{\text{O}}{\longrightarrow}$   $\stackrel{\text{O}}{\longrightarrow}$   $\stackrel{\text{N}}{\longrightarrow}$   $\stackrel{\text{N}}{\longrightarrow}$   $\stackrel{\text{O}}{\longrightarrow}$   $\stackrel{\text{N}}{\longrightarrow}$   $\stackrel{\text$ 

#### 09/847654

RN 119051-51-3 HCAPLUS

.alpha.-Thymidine, 5'-O-[[[5-[(6-chloro-2-methoxy-9-acridinyl)amino]pentyl]oxy]hydroxyphosphinyl]-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-, 3'-(dihydrogen phosphorothioate) (9CI) (CA INDEX NAME)

PAGE 1-A

Me 
$$\stackrel{\text{H}}{\underset{\text{N}}{\longrightarrow}}$$
  $\stackrel{\text{O}}{\underset{\text{O}}{\longrightarrow}}$   $\stackrel{\text{CH}_2-\text{O}}{\underset{\text{O}}{\longrightarrow}}$   $\stackrel{\text{CH}_2-\text{O}}{\underset{\text{O}}{\longrightarrow}}$   $\stackrel{\text{O}}{\underset{\text{O}}{\longrightarrow}}$   $\stackrel{\text{CH}_2-\text{O}}{\underset{\text{O}}{\longrightarrow}}$   $\stackrel{\text{O}}{\underset{\text{O}}{\longrightarrow}}$   $\stackrel{\text{CH}_2-\text{O}}{\underset{\text{O}}{\longrightarrow}}$   $\stackrel{\text{O}}{\underset{\text{O}}{\longrightarrow}}$   $\stackrel{\text{O}}{\underset{\text{O}}}$   $\stackrel{\text{O}}{\underset{\text{O}}{\longrightarrow}}$   $\stackrel{\text{O}}{\underset{\text{O}}{\longrightarrow}}$   $\stackrel{\text{O}}{\underset{\text{O}}}$   $\stackrel{\text{O}}{\underset{\text{O}}}$   $\stackrel{\text{O}}{\underset{\text{O}}}$   $\stackrel{\text{O}}{\underset{\text{O}}}$   $\stackrel{\text{O}}{\underset{\text{O}}}$   $\stackrel{\text$ 

PAGE 2-A

Cl

PAGE 2-B

PAGE 3-A

RN 119051-52-4 HCAPLUS
.alpha.-Thymidine, 5'-O-[[[5-[(6-chloro-2-methoxy-9-acridinyl)amino]pentyl]oxy]hydroxyphosphinyl]-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-, 3'-[S-[2-(4-azidophenyl)-2-oxoethyl] hydrogen phosphorothioate] (9CI) (CA INDEX NAME)

PAGE 1-B

Searcher :

Shears

308-4994

PAGE 2-B

PAGE 2-A

C1

PAGE 3-A

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RN 119051-53-5 HCAPLUS

.alpha.-Thymidine, 5'-O-[[[5-[(6-chloro-2-methoxy-9-acridinyl)amino]pentyl]oxy]hydroxyphosphinyl]-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-, 3'-[S-(carboxymethyl) hydrogen phosphorothioate]
(9CI) (CA INDEX NAME)
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Absolute stereochemistry.

09/847654

PAGE 1-A

PAGE 1-B

308-4994

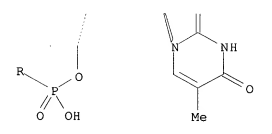
Searcher : Shears

 $\approx_0$ 

09/847654

PAGE 2-A

PAGE 3-A



PAGE 4-A

PAGE 4-B

\_\_OMe

```
RN 119082-35-8 HCAPLUS

cn .alpha.-5'-Cytidylic acid, .alpha.-thymidylyl-(5'.fwdarw.3')-.alpha.-thymidylyl-(5'.fwdarw.3')-.alpha.-thymidylyl-(5'.fwdarw.3')-.alpha.-adenylyl-(5'.fwdarw.3')-2'-deoxy-.alpha.-adenylyl-(5'.fwdarw.3')-.alpha.-thymidylyl-(5'.fwdarw.3')-.alpha.-thymidylyl-(5'.fwdarw.3')-.alpha.-thymidylyl-(5'.fwdarw.3')-2'-deoxy-,5'-[5-[(6-chloro-2-methoxy-9-acridinyl)amino]pentyl] ester (9CI)

(CA INDEX NAME)
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PAGE 2-B

\_\_\_\_ OMe

PAGE 3-B

ΙT 119051-30-8P

> RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, as intermediate for antitumor, antimicrobial , and antiparasitic agents)

RN

119051-30-8 HCAPLUS .alpha.-3'-Thymidylic acid, 5'-O-[[[5-[(6-chloro-2-methoxy-9-CN acridinyl)amino]pentyl]oxy]hydroxyphosphinyl]-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-.alpha.-thymidylyl-(3'.fwdarw.5')-, 3'-(4-chlorophenyl) 3'-(2-cyanoethyl) ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Shears Searcher : 308-4994

PAGE 1-B

PAGE 2-A

Searcher : Shears

308-4994

PAGE 3-A

L11 ANSWER 16 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1988:182738 HCAPLUS

DOCUMENT NUMBER: 108:182738

TITLE: Mechanism of UV endonuclease V cleavage of

abasic sites in DNA determined by carbon-13

labeling

AUTHOR(S): Manoharan, Muthiah; Mazumder, Abhijit; Ransom,

Stephen C.; Gerlt, John A.; Bolton, Philip H. CORPORATE SOURCE: Dep. Chem. Biochem., Univ. Maryland, College

Park, MD, 20742, USA

SOURCE: J. Am. Chem. Soc. (1988), 110(8), 2690-1

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal LANGUAGE: English

The synthetic heptameric heteroduplex of d(GCGDGCG) paired with d(CGCACGC), where D is the deoxyribose abasic site, was a substrate for UV-endonuclease (endonuclease V) from phage T4. The structure of the product contg. the carbohydrate derived from the abasic site was established using a heteroduplex in which the abasic site was labeled with 13C in the 1- and 3-C atoms. The 13C NMR spectra of the endonuclease-catalyzed reaction as a function of time revealed that the endonuclease catalyzes a .beta.-elimination reaction to yield an .alpha.,.beta.-unsatd. aldehyde; this product subsequently undergoes the slow addn. of nucleophiles present in the reaction soln. to yield mixts. of 3-C adducts.

IT **112969-11-6** 

RL: RCT (Reactant)

(cleavage of, by endonuclease V of phage T4, mechanism of)

RN 112969-11-6 HCAPLUS

CN Guanosine, 2'-deoxyguanylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5)-2-deoxy-D-erythro-pentos-3-O-ylphosphinico-(3.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-

(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-

(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-

(3'.fwdarw.5')-2'-deoxycytidine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 112969-10-5 CMF C63 H82 N26 O40 P6

## PAGE 1-A

$$\begin{array}{c|c}
OH & & & & & H & & \\
-O-P-O-CH_2 & & & & & & H & \\
\hline
O & & & & & & \\
O & & & & & & \\
O & & & & & & \\
O & & & & \\
O & & & \\$$

PAGE 2-A

PAGE 2-B

CM 2

CRN 111350-51-7

CMF C66 H85 N27 O39 P6

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

$$H_2N$$
 $N$ 
 $R$ 
 $R$ 
 $R$ 
 $O$ 
 $OH$ 

L11 ANSWER 17 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1980:111255 HCAPLUS

DOCUMENT NUMBER: 92:111255

TITLE: Synthesis of a promoter region of

bacteriophage fd DNA. I. Chemical
synthesis of oligodeoxyribonucleotides

corresponding to the 5'-terminal fragment of the

"minus" strand of the promoter

AUTHOR(S): Efimov, V. A.; Chakhmakhcheva, O. G.

CORPORATE SOURCE: M. M. Shemyakin Inst. Bioorg. Chem., Moscow,

USSR

SOURCE: Bioorg. Khim. (1979), 5(9), 1329-40

CODEN: BIKHD7

DOCUMENT TYPE: Journal

AB Deoxyribonucleotides d(A-A-A-T-C-A-G-G-T-C-T-T) and d(A-C-C-C-T-G-T-C-T-A) corresponding to the (+15)-(-8) fragment of the minus strand of the G2 promoter region of phage fd DNA were synthesized by the phosphodiester method according to 1+1+1+2+2+2+4

and 1+1+1+2+2+3 schemes. The obtained compds. were

5'-phosphorylated by [.gamma.-32P] and T4 polynucleotide kinase and their structures confirmed by nucleotide mapping techniques.

IT 58781-33-2

RL: RCT (Reactant)

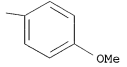
(nucleotide coupling of, with thymidine dinucleotide)

RN 58781-33-2 HCAPLUS

CN Thymidine, 5'-0-[(2-cyanoethoxy)hydroxyphosphinyl]-2'-deoxy-N-(4-methoxybenzoyl)cytidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A



L11 ANSWER 18 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1978:148031 HCAPLUS

DOCUMENT NUMBER:

CORPORATE SOURCE:

88:148031

TITLE:

T4 polynucleotide ligase catalyzed joining of

short synthetic DNA duplexes at base-paired ends

AUTHOR(S):

Deugau, Ken V.; Van de Sande, Johan H.

Div. Med. Biochem., Univ. Calgary, Calgary,

Alberta, Can.

SOURCE:

Biochemistry (1978), 17(4), 723-9 CODEN: BICHAW; ISSN: 0006-2960

DOCUMENT TYPE:

Journal

LANGUAGE:

English

The self-complementary octanucleotide, dT-A-G-T-A-C-T-A, was synthesized and its sequence confirmed by 2-dimensional fingerprinting. Under conditions used for the phage T4 polynucleotide ligase reaction, this oligonucleotide forms a dimeric duplex which shows a Tm of 18.degree.. The optimal rate of joining of the 32P-labeled duplex occurs between 12 and 15.degree.. rate is highly concn.-dependent, as expected for a bimol. process. Polyacrylamide gel electrophoretic anal. of this reaction shows the presence of products up to 120 nucleotides in length. In a denaturing gel, each product appears as double band due to the presence of its 5'-adenylylated activated intermediate. Substrates >8 base pairs are utilized more rapidly than the 8 base pair duplex, indicating that the T4 ligase has a higher affinity for longer substrates. The low level of nicked intermediates suggests that the joining of both strands requires 2 steps, the rates of which must be similar.

IT 58781-33-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and coupling reaction of)

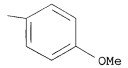
RN 58781-33-2 HCAPLUS

CN Thymidine, 5'-0-[(2-cyanoethoxy)hydroxyphosphinyl]-2'-deoxy-N-(4-methoxybenzoyl)cytidylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B



LANGUAGE:

L11 ANSWER 19 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1970:425810 HCAPLUS

DOCUMENT NUMBER: 73:25810

TITLE: Oligonucleotidic compounds. XXXVI. Synthesis

of uridylyl-(5'.far.3)-uridylyl-(5'.far.5')uridylyl-(3'.far.5')-uridine and its priming activity for polynucleotide phosphorylase

activity for polynucieotide phosph

AUTHOR(S): Smrt, Jiri; Cramer, Friedrich CORPORATE SOURCE: Cesk. Akad. Ved., Prague, Czec.

CORPORATE SOURCE: Cesk. Akad. Ved., Prague, Czech. SOURCE: Collect. Czech. Chem. Commun. (1970), 35(5),

1456-63

English

CODEN: CCCCAK

DOCUMENT TYPE: Journal

AB The title compd. (I) was obtained in 7% yield by condensation of

2'-O-tetrahydropyranyluridylyl-(3' .fwdarw. 5')-2',3'-O-ethoxymethyleneuridine (II) and 5'-O-phosphono-2'-O-tetrahydropyranyluridylyl-(3' .fwdarw. 5')-2',3'-O-

ethoxymethyleneuridine (III) in the presence of 2,4,6-iso-Pr3C6H2SO2Cl. II was prepd. from 2'-O-tetrahydropyranyl-5'-Oacetyluridine 3'-phosphate and 2',3'-O-ethoxymethyleneuridine by the N, N'-dicyclohexylcarbodiimide method, and III was obtained by phosphorylation of II. The priming activity of I, estd. in the polymn. of ADP by primer-dependent polynucleotide phosphorylase (IV), was almost double that of uridylyl-(3' .fwdarw. 5')-uridine. Participation of both portions of the mol. contg. internucleotide linkage in the interaction with the enzyme accounted for this increased activity. I also primed the reaction of IDP and IV in the presence of ribonuclease T1 yielding 3'-IMP and 3'-Ophosphorylinosinyl-(5' .fwdarw. 3')-uridylyl-(5' .fwdarw. 3')-uridylyl-(5' .fwdarw. 5')-uridylyl-(3' .fwdarw. 5')-uridylyl-(4' .fwdarw. 5')-inosine 3'-phosphate, which was degraded by bacterial alk. phosphatase and pancreatic ribonuclease to yield 3'-0-phosphoryluridylyl-(5' .fwdarw. 5')-uridine 3'-phosphate, inosine, and 3'-UMP approx. in the ratio 1:2:2.

IT 28440-24-6P

RN 28440-24-6 HCAPLUS

CN Uridine, uridylyl-(5'.fwdarw.3')-uridylyl-(5'.fwdarw.5')-uridylyl-(3'.fwdarw.5')-(8CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

L11 ANSWER 20 OF 20 HCAPLUS COPYRIGHT 2002 ACS

1969:422298 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 71:22298

TITLE: Polynucleotides. XCII. Synthesis of a

deoxyribododecanucleotide containing specific

amino acid codons

Kumar, Ashok; Khorana, Har G. AUTHOR(S):

Univ. of Wisconsin, Madison, Wis., USA CORPORATE SOURCE: SOURCE: J. Amer. Chem. Soc. (1969), 91(10), 2743-9

CODEN: JACSAT

DOCUMENT TYPE: Journal English LANGUAGE:

High mol. wt. deoxyribopolynucleotides with repeating nucleotide sequences were previously shown to direct the synthesis of polypeptides in the bacterial cell-free protein synthesizing system. With the aim of synthesizing polypeptides contg. specific amino acid sequences via nucleic acid templates, a deoxyribododecanu-cleotide was now synthesized. The synthetic polynucleotide, d-A-T-G-C-A-C-T-C-T-T-A-G, contains at the appropriate 5' end the trinucleotide sequence A-T-G, which stands for formylmethionine and initiates the synthesis of peptide chain, and at the 3' end the sequence T-A-G, which should terminate and release the polypeptide chain. The codons selected for internal positions were C-A-C (histidine) and T-C-T (serine). The protected trinucleotide blocks d-MMTr-ABZpTpGiso-Bu, d-pCAnpABzpCAn-OAc, d-pTpCAnpT-OAc, and d-pTpABzpGiso-Bu-OAc were prepd. by stepwise methods using the protected nucleoside and nucleotides. The blockwise condensations of the protected trinucleotides to form the dodecanucleotide were carried out using mesitylenesulfonyl chloride as the condensing agent, yields in the individual steps being 30-50%.

ΙT 24638-85-5P 24816-20-4P 24934-67-6P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of) 24638-85-5 HCAPLUS RN

Adenosine, 5'-O-[(2-cyanoethoxy)hydroxyphosphinyl]-thymidylyl-(3'.fwdarw.5')-N-benzoyl-2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 24816-20-4 HCAPLUS

CN Cytidine, P-(2-cyanoethyl)-5'-O-phosphonothymidylyl-(3'.fwdarw.5')-2'-deoxy-N-(4-methoxybenzoyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

RN 24934-67-6 HCAPLUS

CN Adenosine, 5'-O-[(2-cyanoethoxy)hydroxyphosphinyl]-2'-deoxy-N-(4-methoxybenzoyl)cytidylyl-(3'.fwdarw.5')-N-benzoyl-2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

ELLE REGISTRY ENTERED AT 15:32:52 ON 09 OCT 2002 52-SEA\_FILE=REGISTRY ABB=ON PLU=ON (112591-87-4/BI OR 119051-51-3/BI OR 119082-35-8/BI OR 167212-06-8/BI OR 446017-73-8/BI OR 446017-74-9/BI OR 58781-33-2/BI OR 112591-90-9/BI OR 112969-11-6/BI OR 119051-30-8/BI OR 119051-52-4/BI OR 119051-53-5/BI OR 119874-42-9/BI OR 120886-02-4/BI OR 130882-87-0/BI OR 131401-19-9/BI OR 131401-20-2/BI OR 131401-22-4/BI OR 131401-23-5/BI OR 131401-24-6/BI OR 131401-25-7/BI OR 131419-77-7/BI OR 131419-78-8/BI OR 131419-79-9/BI OR 131419-80-2/BI OR 151837-15-9/BI OR 161054-62-2/BI OR 166887-47-4/BI OR 166887-51-0/BI OR 166887-52-1/BI OR 166887-53-2/BI OR 167211-99-6/BI OR 167212-03-5/BI OR 167212-04-6/BI OR 167212-07-9/BI OR 180840-08-8/BI OR 185997-79-9/BI OR 216485-50-6/BI OR 216485-51-7/BI OR 216485-52-8/BI OR 216485-53-9/BI OR 216485-54-0/BI OR 216485-55-1/BI OR

24638-85-5/BI OR 24816-20-4/BI OR 24934-67-6/BI OR 28440-24-6/BI OR 331953-81-2/BI OR 403717-05-5/BI OR 403717-06-6/BI OR 403717-07-7/BI OR 403717-08-8/BI)

FILE CAOLD ENTERED AT 15:33:10 ON 09 OCT 2002 L13 0 S L12

FILE 'USPATFULL' ENTERED AT 15:33:15 ON 09 OCT 2002

L14 ANSWER 1 OF 5 USPATFULL

ACCESSION NUMBER: 2002:55001 USPATFULL

Antimicrobial compounds and methods for their use TITLE:

INVENTOR(S): Dale, Roderic M. K., Wilsonville, OR, UNITED

Gatton, Steven L., Lake Oswego, OR, UNITED STATES

Arrow, Amy, Bethel, ME, UNITED STATES

Thompson, Terry, West Linn, OR, UNITED STATES

NUMBER KIND DATE US 2002032164 20020314 PATENT INFORMATION: A1 APPLICATION INFO.: US 2001-847654 A1 20010503 (9)

Continuation-in-part of Ser. No. US 1999-281858, RELATED APPLN. INFO.:

filed on 31 Mar 1999, PENDING

Continuation-in-part of Ser. No. US 1998-222009,

filed on 30 Dec 1998, GRANTED, Pat. No. US

6211349 Utility APPLICATION

Beth A. Burrous, FOLEY & LARDNER, Washington LEGAL REPRESENTATIVE:

Harbour, 3000 K Street, N.W., Suite 500,

Washington, DC, 20007-5109

NUMBER OF CLAIMS: 36 EXEMPLARY CLAIM: 1

DOCUMENT TYPE:

FILE SEGMENT:

NUMBER OF DRAWINGS: 8 Drawing Page(s)

LINE COUNT: 1406

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides protonated compounds having antimicrobial activity. The invention also provides antimicrobial AB compositions comprising protonated compounds of the invention. The protonated compounds of the invention provide efficacious

antimicrobial activity against resistant strains of bacteria and

opportunistic fungi.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L14 ANSWER 2 OF 5 USPATFULL

ACCESSION NUMBER: 2001:48036 USPATFULL

TITLE: Pulmonary delivery of protonated/acidified

nucleic acids

INVENTOR(S): Dale, Roderic M. K., Wilsonville, OR, United

States

Gatton, Steven L., Lake Oswego, OR, United States

Arrow, Amy, Bethel, ME, United States Oligos Etc. Inc., Wilsonville, OR, United States PATENT ASSIGNEE(S):

(U.S. corporation)

NUMBER KIND DATE \_\_\_\_\_ US 6211162 B1 20010403 US 1999-282824 · 19990331 (9) PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1998-222009, filed on 30 Dec 1998 Utility DOCUMENT TYPE: FILE SEGMENT: Granted PRIMARY EXAMINER: Clark, Deborah J. R.

PRIMARY EXAMINER:
ASSISTANT EXAMINER: Chen, Shin-Lin

LEGAL REPRESENTATIVE: Bozicevic, Field & Francis LLP, DeVore, Dianna L.

NUMBER OF CLAIMS: 1.5 EXEMPLARY CLAIM: 1 LINE COUNT: 1501

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides a method of treating bacterial respiratory infections by pulmonary administration of protonated/acidified nucleic acids. These modified nucleic acids are effective as bactericidal and/or bacteriostatic agents without regard to the class of bacteria, so are especially useful when diagnosis is difficult or when multiple infectious organisms are present. The antibiotic activity of nucleic acids of the invention is not dependent on either the specific sequence of the nucleic acid or the length of the nucleic acid molecule.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L14 ANSWER 3 OF 5 USPATFULL

ACCESSION NUMBER: 1999:128744 USPATFULL

Nucleic acid probes chemically modified at 5'(OH) TITLE:

and/or at 3'(OH) for the purpose of introducing one or more non-radioactive marking elements at these sites, and method for preparing the same

INVENTOR(S): De Vos, Marie-Joelle, Feluy, Belgium

Bollen, Alex, Itterbeek, Belgium

La Region Wallone, Brussels, Belgium (non-U.S.

PATENT ASSIGNEE(S):

corporation)

NUMBER KIND DATE US 5969128 WO 9419364 PATENT INFORMATION: 19991019 19940901 APPLICATION INFO.: US 1995-507283 19950821 (8) WO 1994-BE13 19940218 19950821 PCT 371 date 19950821 PCT 102(e) date

NUMBER DATE

PRIORITY INFORMATION: BE 1993-160 19930219

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

Houtteman, Scott W. PRIMARY EXAMINER:

Sughrue, Mion, Zinn, Macpeak & Seas, PLLC LEGAL REPRESENTATIVE: 6

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 15 Drawing Figure(s); 8 Drawing Page(s)

LINE COUNT: 1681

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AΒ The probe comprises: a) an oligonucleotide or oligodeoxyribonucleotide part constituted by a DNA or RNA nucleic acid sequence S, depending on the type of molecule to be detected, and b) a non-nucleotide part possessing chemical properties enabling direct or indirect atttachment of one or more detection units or marking elements M detectable non-isotopically by production of colour or light. The probe is characterized by the fact that part b) is constituted by a chain of phosphate units interspersed with alkyl groups, viz.: b1) certain alkyl groups uniting the different phosphate groups and presenting no special functionality b2) alkyl groups presenting primary amine groups which allow splicing with varied reagents to carry out direct or indirect detection, the b2) groups being bonded to part a) or sequence S by way of groups b1). Sequence S is bonded at its 5' and/or 3' extremity to one or more marking elements M. The probes of this type are used to detect and diagnose hereditary genetic diseases, oncogenes, viral, bacterial or parasitic diseases.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L14 ANSWER 4 OF 5 USPATFULL

ACCESSION NUMBER: 97:14599 USPATFULL

TITLE: Improvements in or relating to DNA cloning

techniques and products for use therewith

INVENTOR(S): Taylor, Philip N., Nr Dartford, United Kingdom

PATENT ASSIGNEE(S): The University of Hull, Hull, United Kingdom

(non-U.S. corporation)

	NUMBER	KIND DATE	
DAMENIM THEODMANION.	UC E CO 4100	19970218	
PATENT INFORMATION:	US 5604122 WO 9319186	19930930	
APPLICATION INFO.:	US 1994-307713	19941114	(8)
	WO 1993-GB584	19930322	
		19941114	PCT 371 date
		19941114	PCT 102(e) date

NUMBER	DATÉ	
3 1992-6210	19920321	

PRIORITY INFORMATION: GB 1992-6210 DOCUMENT TYPE: Utility

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Fleisher, Mindy ASSISTANT EXAMINER: Brusca, John S.

LEGAL REPRESENTATIVE: Leydig, Voit & Mayer, Ltd.

NUMBER OF CLAIMS: 17
EXEMPLARY CLAIM: 1
LINE COUNT: 873

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A method of cloning foreign DNA into a DNA vector comprising ligating:

- (1) a DNA vector having a single stranded DNA overhang at each end, said overhangs being mutually incompatible so as to prevent self-religation, with
- (2) a linear piece of foreign DNA having a single stranded DNA

overhang at each end,

each foreign DNA overhang being complementary to but at least one base shorter than each of the vector overhangs and being capable of base pairing along the entire length of the overhang with one of the vector overhangs, and sealing the gap by either transforming the double stranded DNA having a gap therein into a suitable bacterium or transfecting it into a suitable bacterium after packaging it into a suitable bacteriophage.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L14 ANSWER 5 OF 5 USPATFULL

96:113828 USPATFULL ACCESSION NUMBER:

Method of cleaving specific strands of RNA TITLE:

INVENTOR(S): Torrence, Paul, Silver Spring, MD, United States

Silverman, Robert, Shaker Heights, OH, United

States

Maitra, Ratan, Euclid, OH, United States

Lesiak, Krystyna, Gaithersburg, MD, United States

The Cleveland Clinic Foundation and National PATENT ASSIGNEE(S):

Institutes of Health, Bethesda, MD, United States

(U.S. corporation)

NUMBER KIND DATE \_\_\_\_\_\_\_

PATENT INFORMATION: US 5583032 19961210 US 1993-123449 19930917 (8)

APPLICATION INFO.:

Continuation-in-part of Ser. No. US 1992-965666, RELATED APPLN. INFO.:

filed on 21 Oct 1992, now abandoned

DOCUMENT TYPE: Utility

FILE SEGMENT: Granted

PRIMARY EXAMINER: Rories, Charles C. P. LEGAL REPRESENTATIVE: Pennie & Edmonds

NUMBER OF CLAIMS: 21 EXEMPLARY CLAIM: 1

12 Drawing Figure(s); 12 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 2560

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A method of using a chimeric molecule made up of an antisense oligonucleotide attached to a 2',5'-oligoadenylate molecule to specifically cleave a sense strand of RNA, wherein the antisense oligonucleotide of the chimeric molecule is hybridized to the sense strand of RNA in the presence of 2',5'-dependent RNase.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

FILE 'HOME' ENTERED AT 15:33:29 ON 09 OCT 2002

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